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PEPTIDE FRAGMENTS OF MURINE EPIDERMAL GROWTH FACTOR AS LAMININ RECEPTOR TARGETS

2 This invention delates to the use of (synthetic and 3 modified) lamenin receptor-targetted ligands for the 4 treatment of any genic diseases such as proliferative 5 retinopathies and metastatic cancer as well as for the 6 treatment of Candida spp. infections, or parastic 7 infestations such as leishmania and trichomonas 8 9 vaginalis. 10 Laminin antagonists (which are anti-angiogenic) can be 11 used to inhibit secondary turgur spread (by inhibiting 12 tumour cell attachment) and to grevent growth of 13 metastatic secondaries (by inhibiting 14

neovascularisation).

diabetic retinopathy).

Laminin agonists (which promote angiogenesis) dould be used to treat retinopathy of prematurity, and could also be used to promote wound healing (for example in corneal epithelium).

used to treat other angiogenic disorders (such as

These antagonists could also be

Both the antagonists and the agonists would be expected to inhibit parasite binding to tissue surfaces and would thus prevent infection or infestation.

Angiogenic diseases are those disorders which are directly caused by, or complicated by the inappropriate growth of new blood vessels. The major angiogenic diseases include the common metastatic solid tissue

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WO 99/54356 PCT/GB99/01211

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1	cancers	(breast,	gastrointestinal,	lung,	prostatic,
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- 2 etc), diabetic retinopathy, neovascular glaucoma,
- 3 rheumatoid arthritis and psoriasis. Angiogenesis is
- 4 the rate-limiting step in the growth of secondary
- 5 tumours; inhibition of their neovascularisation is
- 6 known to stop their growth.

8 In this field it is already known that the native

- 9 ligand of the 67kDa laminin receptor (67LR) is
- 10 encompassed by the linear sequence of amino acids 925-
- 11 933 of the laminin β -1 (previously known as laminin B1
- or b1) chain (numbering refers to the mature murine
- laminin β -1). Synthetic laminin β -1₉₂₅₋₉₃₃ (single letter
- amino acid code: CDPGYIGSR-NH2) has been shown to
- inhibit tumour establishment in mice, by inhibiting
- 16 attachment of tumour cells to basement membranes. It
- has also been demonstrated that laminin β -1₉₂₅₋₉₃;
- inhibits angiogenesis in the chick.

20 However, synthetic laminin-derived peptide (laminin

- 21 β -1₉₂₅₋₉₃₃) stimulates angiogenic events in mammalian
- cells (in which it acts as a pure 67LR agonist), making
- 23 it useless as the basis of a human therapy.

25 It is one object of the present invention to provide a

26 medicament to treat angiogenic diseases.

28 The present invention provides a peptide factor derived

- from murine epidermal growth factor (EGF) peptide for
- 30 use in the preparation of a medicament for the
- 31 treatment of angiogenic diseases.
- 33 The mechanism by which EGF derived peptides inhibit new
- 34 blood vessel formation is through their antagonism of
- 35 the high affinity 67 kDa laminin receptor (67LR) found
- 36 on endothelial cells.

PCT/GB99/01211 WO 99/54356

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1 '	The	peptides	have	the	additional	effect	of	inhibiting
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- tumour cell attachment to basement membranes, and may 2
- be used to prevent solid cancer spread in cases where 3
- cancer cells have been identified circulating in the 4
- 5 blood.

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- Modified peptides may be protected from proteolytic 7
- degradation by substitution of key residues with 8
- unnatural amino acid analogues at susceptible bonds, 9
- such as tyrosine analogues (at position 5) and arginine 10
- analogues (at position 9). The peptides may be capped 11
- at N- and C-termini (with acetyl and amide groups 12
- respectively) and at the thiol groups of the cysteines 13
- (with acetamido methyl groups). 14

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- Typically the peptide is an antagonist of the 67kDa 16
- Laminin Receptor (67LR). 17

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- The peptide factor is based on amino acid residues 33 19
- to 42 of murine epidermal growth factor (mEGF). 20

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The amino acid sequence of mEGF- (33-42) is CVIGYSGDRC. 22

23

- Preferably the sequence of peptide factor is modified 24
- from the natural sequence to protect the peptides from 25
- protease attack. 26

27

- Preferred substitutions include the use of tyrosine 28
- analogues at position 5 and arginine analogues at 29
- position 9. 30

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- Preferably the peptide factor is capped at the N 32
- terminal with an acetyl group. 33

34

- Preferably the peptide factor is capped at the C 35
- terminal with an amide group. 36

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WO 99/54356 PCT/GB99/01211

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Preferably the thiol groups of cysteines are capped with acetamido methyl groups.

3

In one embodiment the synthetic peptide has the sequence

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Acetyl-C-[S-Acm]-VIGYSGDR-C-[S-Acm]NH2

8

A preferred tyrosine analogue is Tic-OH.

10 11

A preferred arginine analogue is Citrulline.

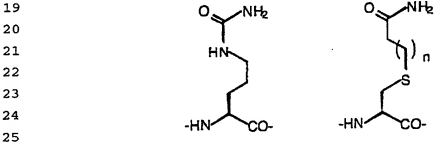
12

13 The structure of Citrulline and other potential 14 arginine analogues are shown below.

15 16

Citrulline and analogues

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citrulline cysteine-derived analogues
{prepared by reaction of cysteine with Br-(CH₂)_n-CONH₂}

28 29

> 35 36

30 S NH₂
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thiono-citrulline

homo-glutamine

{prepared by reaction of ornithine with ammonium isothiocyanate}

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Preferably the peptide is truncated to a shorter 1

peptide without losing its antagonistic character. 2

3

The invention further provides a peptide agonist. 4

5

The agonist may be the native sequence (single letter 6

7 amino acid code: CDPGYIGSR-NH2) or may have the tyrosine

substituted by any of a variety of tyrosine analogues 8

such as the comformationally restricted Tic-OH or 9

2',6'-dimethyl-beta-methyl-tyrosines, 2-0-methyl and 2-10

O-ethyl-tyrosine and the like. 11

12

The agonist may be useful in healing endothelial cell 13

wounding. 14

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For example, corneal endothelial cells can be damaged 16

during cataract operations and this damage does not 17

self-repair because these endothelial cells do not 18

divide. Healing can only be effected by cell migration 19

and spreading, and this may be promoted by the agonist. 20

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24

In order to explore possible conformations for the 22

parent mEGF33-42 peptide, it was modelled using molecular 23

dynamics. Based on these conformations a strategy has

been predicted to provide proteolytic protection by 25

being able to identify residues that are important to 26

27 the maintenance of a three-dimensional conformation

essential for 67LR recognition. 28

29

The following is a description of some examples of 30

modifications and uses of the invention. 31

32

On the basis of the modelled structures, it was 33 1. found that the arginine residue participated in H-34 bonding, and speculated that this charge may not 35

be important. A peptide was synthesised based on 36

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WO 99/54356 PCT/GB99/01211 6 1 mEGF33-42, in which the arginine residue at position 2 41 was replaced by citrulline (an uncharged arginine mimetic with similar H-bonding 3 potential). This peptide provided to act as a 4 more potent 67LR antagonist and was found to be 5 resistant to trypsin degradation. 6 7 Double substitution of tyrosine, with Tic-OH and 2. 8 arginine, with citrulline, to produce a mEGF₃₃₋₄₂-9 10 derived peptide resistant to both chymotrypsinlike and trypsin-like proteases. 11 12 13 3. Replacement of susceptible peptide bonds in mEGF33. 4, with protease-resistant peptide bond isosteres 14 (such as thionopeptide or methylene amino bonds). 15 16 Conformationally restricted analogues may give 17 4. improved potency due to the essential 3-18 dimensional conformation being stabilised. For 19 example, it should be possible to increase the 20 rigidity of the molecule by replacing each of the 21 central glycine residues in turn by α, α -dialkyl 22 substituted amino acids such as α -amino isobutyric 23 acid (AIB) or aminocyclopropane carboxylic acid 24 (ACPCA). Alternatively, the helical turn (which 25 we have identified as essential) could be 26 stabilised by bridging with suitable intra-chain 27 28 linkers, such as a disulphide bond between N- and C-terminal [D] or [L]-cysteines. 29 30 31 EXAMPLE 1 32 33 The invention is demonstrated with reference to 34 the following figures wherein. 35

Figure 1a depicts a flat mount retina showing the

effects of ROP and Figure 1b depicts a retina from laminin-agonist treated mouse showing re-

3 canalisation of vessels.

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Treatment of Retinopathy of Prematurity (ROP)

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Severely premature babies are at risk of developing retinopathies due to their being exposed to high oxygen levels post-partum. life-saving intervention compensates for poor lung development but has the unfortunate side-effect of causing unnaturally hyperoxic conditions in the The direct effect of this is to remove the normal hypoxic cues for endothelial migration, resulting in inhibition of capillary growth and vaso-obliteration. When these babies are returned to room air, hypoxic stimuli are restored and retinal angiogenesis is again induced. However, the newly induced angiogenesis is chaotic and uncontrolled, often resulting in abnormal penetration of vessels into the vitreous (see Figure 1a, below). It is the uncontrolled growth of these blood vessels that ultimately leads to loss of visual activity.

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It has now been shown that laminin agonist treatment can reverse the effects of both hyper-oxic induced vaso-ablation as well as norm-oxic-induced angiogenesis in a murine model of retinopathy of prematurity (ROP). In this model, development of ROP can be prevented by treatment of neonates with daily injections (intraperitoneal) of $10\mu g$ of synthetic laminin β - $1_{925-933}$ (also referred to as laminin $B1_{925-933}$, single letter amino acid code:CDPGYIGSR-NH₂). See Figure

WO 99/54356

PCT/GB99/01211

1b in comparison with 1a. Treatment with laminin 1 agonist (Figure 1b) prevents the uncontrolled 2 angiogenic response of ROP (Figure 1a) and 3 promotes re-canalisation of areas of vaso-4 obliteration. 5 6 The invention is demonstrated with reference to 7 the following figures wherein Figure 1a depicts a 8 flat mount retina showing the effects of ROP 9 10 Figure 1b depicts a retina from laminin-agonist 11 treated mouse showing re-canalisation of vessels. 12 13 Murine model of proliferative retinopathy 14 15 Litters of 7 day old C57-BL/6J mice, together with 16 their nursing dams, are exposed to 80% oxygen in 17 an incubator maintained at 23°C and with a gas 18 exchange of 1.5L/min for 5 days according to the 19 protocol described by Stitt et al. (1998). 20 postnatal day 12 (P12) the animals are returned to 21 room air and sacrificed at various times post-22 hyperoxia. Animals are treated with daily i.p. 23 injections of either laminin agonists (10µg per 24 head per day) or vehicle control. Groups of room 25 air controls are maintained in parallel with 26 hyperoxia-exposed animals. Home Office project 27 and personal licenses are held for this work. All 28 animals are housed and maintained in accordance 29 with the ARVO regulations for animal care in 30 research. 31 32 Animals are sacrificed at pre-determined key 33 34 stages in the vaso-obliteration (P7-P12), ischaemia (P12 onwards) and vaso-proliferative 35 responses (P12-21). At sacrifice, terminally 36

anaesthetised animals have a single eye enucleated and the retina removed to be snap-frozen for later RNA-extraction (see below). The fellow eye is either perfused with fluorescein dextran or enucleated and fixed in 4% paraformaldehyde for histology, immunohistochemistry and in situ hybridisation.

ALTERNATIVE USES

1. Treatment of corneal wounds

The cornea is a delicate transparent structure. Being avascular, corneal wound healing depends upon local self-renewal of the corneal epithelium. This, in turn, depends upon the presence of a mitogenically functional stem cell population ('limbal cells'), which produce replacement cells that migrate and desquamate at the denuded area. Damage to these underlying stem cell populations causes inappropriate re-epithelialization by conjunctival cells followed by matrix deposition and scar formation. The damaging agent may be corrosive chemical or heat burns, erosion by contact lenses, Stevens Johnson disease.

It is known that transplantation of limbal cell autografts from the unaffected eye can restore a stable healing of the corneal epithelium (Kenyon et al., 1996). It has been proposed that harvesting small samples of limbal stem cells, followed by serial culture in vitro would provide greater chance of success (particularly when both eyes are affected) De Luca, et al., 1997). However, with both protocols, correct uptake and controlled migration of these grafted cells into

the corneal epithelium has not been optimised.

We propose that laminin agonists could be used to stimulate the migratory response of the cells prior to grafting, or alternatively topical application of laminin agonists to the wound site could be used to direct migration of the grafted cells to the correct (denuded) area of the cornea.

2. Some microbial pathogens such as Candida albicans, express 67LR and use this as a means of attaching to human basement membranes. It is conceivable that such infections could be abolished by treatment with mEGF₃₃₋₄₂-derived peptides, which would prevent the microbes from adhering to the host.

EXAMPLE 2

Peptide Study

The purpose of the investigation was to determine the molecular target of $mEGF_{(33-42)}$ and to identify the amino acids that are essential for receptor recognition. In addition, the key residues which confer laminin antagonism on $mEGF_{(33-42)}$ were examined.

Two lead compounds were investigated; synthetic laminin β -1 sequence CDPGYIGSR-NH₂ and mEGF₍₃₁₋₄₂₎ sequence AcC(Acm)-VIGYSGDRC-(Acm)-NH₂. Bearing in mind the pure antagonism of the murine EGF peptide, the aims of this study were to identify the key residues responsible for these contrasting activities using alanine scanning, in the context of developing anti-angiogenic drugs for retinopathy treatment.



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		•	_		_			

In addition, using residue	exchange	between	the	two
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- peptides and molecular modelling to predict three-2
- dimensional structure, we wished to further investigate 3
- the role of individual mEGF(13.42) residues in laminin 4
- antagonism. A logical series of peptides was 5
- 6 synthesised and screened for receptor interaction, cell
- adhesion and motility properties (Table 1a and 1b). 7

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MATERIALS AND METHODS

10 11

Peptide synthesis

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13 Peptide sequences based on and mEGF(33-42) were

synthesised on a model 432A peptide synthesizer 14

(Applied Biosystems, Warrington, UK), using standard 15

solid-phase Fmoc procedure (Fields 1990). Synthesis of 16

17 the peptides required successive additions of

derivatized amino acids to form a linear product. 18

19 20

22

Peptides were purified after synthesis using reverse

phase HPLC and purity confirmed by automated amino acid 21

analysis and electrospray mass spectrometry. All

peptide sequences were stored in the presence of 23

desiccant at -20°C until required for biological assay. 24

25

Laminin receptor antibody production

26 27 28

a. Preparation of MAPs

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30 The peptide sequence (PTEDWSAQPATEDWSAAPTA),

corresponding to the COOH-terminal end of the human 31

laminin receptor, was used as the antigen template. 32

Derivation of the peptide, based on a CN-Br cleavage 33

34 fragment of the cDNA sequence encoding human laminin

receptor, has been described elsewhere (Wewer et al 35

36 1986). The antigen was synthesised as an octomeric

WO 99/54356

PCT/GB99/01211

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peptide derivative (MAPs) using automated Fmoc

procedure (Tam 1988).

3 4

5 Table la: Peptide substitution

mEGF ₍₃₃₋	acetyl	ACM Cys	Val	lie	Gly	Тут	Ser	Gly	Asp	Arg	ACM Cys-NH ₂
1	acetyl	ACM Cys	Val	Ite	Gly	Туг	lie	Gly	Asp	Arg	ACM Cys-NH2
TI .	acetyl	ACM Cys	Val	Ile	Gly	Tyr	Ser	Gly	Ser	Arg	ACM Cys-NH ₂
III	acetyl	ACM Cys	Val	Ile	Gly	Tyr	Ile	Gly	Ser	Arg	ACM Cys-NH ₂
TV	acetyl	ACM Cys	Val	lle	Gly	Tyr	Ser	Gly	Asp	Cit	ACM Cys-NH ₂
V	acetyl	ACM Cys	Val	lle	Gly	OH Tie	Ser	Gly	Asp	Arg	ACM Cys-NH ₂

Table 1b: Peptide substitution (alanine scanning)

mEGF ₍₃₃₋	acetyl	ACM Cys	Val	Ile	Gly	Туг	Ser	Gly	Asp	Arg	ACM Cys- NH ₂
VI	acetyl	ACM Cys	Val	Ala	Gly	Тут	Ser	Gly	Asp	Arg	ACM Cys- NH ₂
VII	acetyl	ACM Cys	Ala	Ile	Gly	Тут	Ser	Gly	Asp	Arg	ACM Cys- NH ₂
VIII	acetyl	Ala	Val	ile	Gly	Тут	Ser	Gly	Asp	Arg	ACM Cys- NH ₂
ιX	acetyl	ACM Cys	Val	Ile	Gly	Tyr	Ser 、	Gly	Asp	Arg	Ala- NH ₂
X	acetyl	ACM Cys	Val	Ile	Gly	Tyr	Ala	Gly	Asp	Arg	ACM Cys- NH2

13

b. Immunisation schedule

2

A pre-immune test bleed (5ml) was obtained from the 3 marginal ear vein of a male New Zealand White rabbit 4 (3.2 kg). The bleed was allowed to clot for 2 h at room 5 temperature after which its edge was detached from the 6 wall of the collection vessel. The clot was then 7 allowed to contract overnight at 4°C. Serum was then 8 removed and the residual material pelleted out by 9 centrifugation (10 min at 2,500 g). Extracted serum 10

11 (3.5 ml) was then frozen at -20°C until required.

12

Immunogen was prepared by the emulsion of MAPs (0.5 g 13 antigen in 0.5 ml PBS) in an equivalent volume of 14 adjuvant (Alum Imject; Pierce, Chester, UK). The 15 animals immune system was primed by introducing 16 immunogen (50 μ g) through subcutaneous injection at 17 different sites on the animals back. The rabbit was 18 boosted by both subcutaneous and intramuscular 19 injection, 21 days after priming, using an increased 20 dose of immunogen (800 μ g). Subsequent boosts were 21 performed by intramuscular injection after a further 14 22 days (800 µg immunogen), and thereafter at 21 day 23 intervals. Test bleeds were taken 2 days after each 24 boost and the serum extracted as described above. The 25 animal was boosted and bled a total of three times.

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c. Enzyme-linked immunoabsorbent assay

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ELISA was used to determine the specificity of the antibody prepared against the synthetic MAPs peptide and to determine the efficacy of binding with respect to that of the linear precursor.

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35 Peptides were dissolved in distilled water and diluted

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WO 99/54356 PCT/GB99/01211

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1	to 10 $\mu \mathrm{g/ml}$ in coating buffer. Aliquots (100 $\mu \mathrm{l}$) of
2	either linear or MAPs peptide were then added to the
3	wells of microtitre plates (Microtest III; Becton
4	Dickinson Ltd., Oxford, UK) and incubated overnight at
5	37°C. The wells were then rinsed with 100 μ l wash
6	buffer and air dried. Excess adsorption sites were
7	blocked (1 h incubation at 22°C) by the addition of 10%
8	casein in PBS (0.1 ml/well). Subsequent to the removal
9	of casein solution by aspiration, wells were again
10	rinsed with wash buffer and air dried.
11	
12	Antisera or pre-immune sera were then serially diluted
13	in PBS and 100 μ l of each incubated in peptide coated
14	wells for 1 h at 37°C. After rinsing (0.1 ml wash
15	buffer), 100 μ l per well of 5 μ g/ml secondary antibody
16	(horse-radish peroxidase-conjugated goat anti-rabbit
17	IgG; Amersham International, Aylesbury, UK) was added
18	to each well and the plates incubated at 37°C for 1 h.
19	
20	Wells were again rinsed with wash buffer and 0.1 ml
21	substrate solution (TMB peroxidase) added to each. The
22	plate was then incubated at 22°C for 30 min and the
23	colour reaction stopped by the addition of $0.5M\ H_2SO_4$
24	(0.1 ml/well). Absorbence was measured at 450 nm on a
25	Titertek Multiscan plate reader.
26	
27	d. Purification of IgG fraction
28	·
29	Anti-laminin receptor antiserum was purified using
30	immobilised protein G-sepharose columns (Pharmacia
31	Biotech, Uppasla, Sweden). The columns were

equilibrated with 20 ml sodium phosphate buffer (pH

a 1 ml aliquot loaded onto the column (flow rate 150

ml/h, fraction size 2.5 ml). After exclusion of the

7.0). Antiserum was diluted 1:4 in the same buffer and

PCT/GB99/01211 WO 99/54356

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unbound fraction, as determined by absorbence at 280nm, 1 the IgG component of the antiserum was eluted with 0.1M 2 glycine-HCl (pH 2.7), into tubes containing 0.1 ml Tris 3 (1M), pH 9.0. The eluted IgG fractions were bulked and 4 stored at -20°C until required. 5 б 7 Maintenance of cell cultures 8 Cancer and endothelial cells were maintained in either 9 DMEM (T47-D) or RPMI (SK HEP-1) media, supplemented 10 with 10% FCS, 100 IU/ml penicillin and 100 $\mu g/ml$ 11 streptomycin. Cells were incubated at 37°C in a 12 humidified atmosphere of 95% air: 5% CO2 and media 13 refreshed as required. Cultures (at 80-85% confluence) 14 were routinely passed on removal from monolayer by the 15 action of trypsin (0.25%) and EDTA (0.02%) in CFS. 16 17 The viability of cell populations following 18 trypsinisation was determined by the trypan blue vital 19 dye exclusion test. Populations confirmed as being in 20 excess of 95% viable were used in all studies. 21 22 Media were screened for possible bacterial or fungal 23 contamination by incubating 1ml aliquots with both 24 nutrient and Saboraud dextrose broths (Oxoid Ltd., 25 Basingstoke, UK). Cell populations were routinely 26 monitored for sub-clinical infections by periodically 27 culturing in the absence of antibiotics. 28 Both cell lines and media were examined for the

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presence of contaminating Mycoplasma spp. by the method 31

of Chen (1977). 32

Determination of cell numbers 33

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Single cell suspensions were quantified using an 35

PCT/GB99/01211 WO 99/54356

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ı	automated counter (Coulter Electronics, Harpenden, UK).
2	A 1 ml aliquot of cell suspension was diluted 1 in 20
3	in Isoton and 0.5 ml samples counted. The mean of 5
4	counts was taken and the total number of cells
5	determined. Estimates of cell number were confirmed by
6	counting in a haemocytometer.
7	
8	For microtitre end-point assays, cell numbers were
9	estimated from the crystal violet staining index of the
10	cell line (Kanamaru and Yoshida 1989). Briefly, after
11	removal of media from the assay system cells were fixed
12	with formaldehyde (10% in PBS), and washed with
13	distilled H_2O . Aliquots (100 μ l) of crystal violet
14	solution (0.1% in distilled H_2O) were added to each well
15	and the plates allowed to stand for 30 min. Excess
16	stain was removed by rinsing with distilled H_2O (3 x 100
17	μ l). The wells were then air-dried and the remaining
18	crystal violet extracted with 100 μ l acidified
19	methanol. Absorbance at 620 nm was determined using a
20	Titertek Multiscan spectrophotometer.
21	
22	Proliferation assays
23	
24	The effects of synthetic peptides and growth factors on
25	the growth of breast cancer and endothelial cells were
26	determined as detailed.
27	
28	Exponentially growing cells were harvested by
29	trypsinisation, as previously described. After rinsing
30	and resuspending in the relevant culture media
31	(containing 10% FCS), the cells (100 μ l aliquots) were

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(containing 10% FCS), the cells (100 μ l aliquots) were dispensed into 96-well microtitre plates at a population density of 2 x 104 cells/well (6 wells per experimental condition). Cells were the incubated for 24 h at 37°C after which the media was removed and the

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PCT/GB99/01211 WO 99/54356

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wells rinsed with CFS (3 x 100 μ l), to rid the plates 1 of cells in suspension. Media was then replaced with 2 that containing the relevant controls or treatment 3 supplements as detailed in individual experiments. 4

5 6

Cell numbers were evaluated spectrophotometrically at 620 nm, over the period of assay, after fixing with 10% formaldehyde and staining with crystal violet.

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Proliferative responses were analysed using the 10 Wilcoxan Rank test and significant differences at the p11 < 0.05 level, defined. Results of all growth studies 12 were confirmed in at least 3 individual experiments. 13

14 15

Laminin attachment assay

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Non-tissue culture grade 96-well plates, coated with 17 2.5 μ q murine laminin in 50 μ l CFS per well, were air-18 dried overnight at room temperature. Preliminary 19 experiments indicated that cell attachment was 20 concentration dependent; maximal binding occurred at a 21 laminin coating of 2.5 $\mu g/well$. After rinsing with CFS 22 (100 μ l), the plastic was saturated with casein (0.2% 23 in CFS). Plates were incubated at room temperature for 24 45 min then washed extensively with CFS (3 x 100 μ l). 25

26

After removal of culture media, cells were detached 27 from monolayers by the action of EGTA (0.02% in CFS) at 28 37°C. The cells were then centrifuged at 800 g for 2 29 min and the pellet resuspended in DMEM (T-47D) or RPMI 30 (SK HEP-1). 31

32

Cells, at a population density of 10° cells/ml, were 33 then aliquoted (1 ml) into microfuge tubes containing 34 the individual peptide sequences and incubated for 1 h 35

18

at 37°C. The cells (100 μ l aliquots) were then added to

- 2 the pre-coated multi-well plates and incubated for a
- 3 further 60 min. Incubation media were removed and the
- 4 wells washed with CFS (3 x 100 μ l) to rid the plates of
- 5 non-adherent cells.

6

- 7 Attached cell numbers were evaluated
- 8 spectrophotometrically at 620 nm after fixing with 10%
- 9 formaldehyde and staining with crystal violet.

10

11 Attachment to mEGF (33-42)

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- That $mEGF_{(33-42)}$ bound to the 67kDa laminin receptor was
- 14 demonstrated using a biotinylated derivative of the
- peptide (Acetyl-C-[S-Acm]-VIGYSGDR-C-[S-Acm]-K-[N-
- biotin]-amide) and a modification of the above laminin
- 17 attachment assay.

18

- 19 Briefly, 96-well plates were coated with 100 μ l/well
- streptavidin (5 μ g/ml in carbonate buffer pH 9.6) and
- 21 following an overnight incubation at 37°C, wells were
- 22 washed with CFS (3 x 100 μ l) and the plastic blocked
- with casein (0.2% in CFS). The plates were then
- 24 incubated at room temperature for 45 min and washed
- 25 with CFS as previously detailed. Biotinylated mEGF (33-42)
- in CFS was then aliquoted into the wells (0.1 ml of 100
- μ M) and the plates incubated for 3 h at 37°C.

28

- 29 After a further block with 0.2% casein, the wells were
- washed with CFS (3 x 100 μ l aliquots). Plates were kept
- 31 at 4°C and used within 2 h.

32

- 33 Cells were prepared as above and pre-incubated for 1 h
- 34 at 37°C with serial dilutions of anti-laminin receptor
- 35 polyclonal (see below) or anti-EGF (R1) receptor

19

monoclonal antibodies. Subsequent procedures were as detailed for the laminin attachment assay.

3 4

Laminin receptor binding determinations

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a. Radiolabelling of laminin

7

125 I-laminin was prepared using 125 I-labelled sodium 8 iodide (Amersham, UK) and immobilised chloramine-T 9 (Iodobeads; Pierce, Illinois). Prior to use, the beads 10 were washed with 500 μ l phosphate buffer (pH 6.5) to 11 remove excess reagent from the support. These were then 12 allowed to air dry and individual beads added to a 13 solution of carrier free Na¹²⁵I, diluted with iodination 14 buffer (phosphate buffer pH 7.4). The beads were 15

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Laminin (10 μ g in 10 μ l) was then diluted into the iodination buffer and the system incubated at 20°C for 15 min. The solution was then removed from the reaction vessel and excess Na¹²⁵I and unincorporated ¹²⁵I₂ separated from the iodinated protein by gel filtration on a GF-5 exclusion column (Pierce, Illinois). Iodinated laminin fractions were recovered at a specific activity of approximately 1.2 mCi/mg protein (864 Ci/mmol).

26 27

b. Competition binding estimation

allowed to equilibrate for 5 min.

28 29

Near confluent cultures of T47-D or SK HEP-1 cells were removed from monolayer with 0.02% EGTA and passed through a G-25 syringe needle to produce single cell suspensions. Aliquots of each cell type (106 cells/ml) were dispensed into separate Ependorf tubes (1 ml each) and pelleted. The cells were then resuspended in 1 ml

20

ice-cold RPMI (SK HEP-1) or DMEM (T47-D) containing 1 0.1% BSA and either laminin or synthetic peptide at the 2 concentrations indicated. Iodinated laminin was then 3 added to each cell suspension to give a final 125I-4 laminin concentration of 0.1 nM (approximately 50,000 5

cpm). These mixtures were incubated overnight at 4°C. 6

7

The tubes were then microfuged at 10,000 g and the 8 supernatant removed. After washing the pellet with 500 9 μ l CFS, the remaining radioactivity was determined 10 using a gamma radiation counter. Non-specific binding 11 was determined by incubating cells with a 1000-fold 12 molar excess of unlabelled laminin. All estimations 13 were carried out in triplicate. 14

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 IC_{50} (concentration of unlabelled peptide required to produce 50% inhibition of radioligand binding) and EC50 (effective concentration for 50% inhibition of cell attachment) values were calculated using the Grafit curve-fitting programme (Erithacus Software, London, UK).

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Migration assays

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The method used was basically as described by Albrecht-25 Buehler (1977). Briefly, coverslips (22 x 22 mm) were 26 treated in 5% detergent (7X; ICN Biomedicals) and 27 washed in alcohol to remove grease. After drying, they 28 were immersed in gelatin solution (Sigma, 300 Bloom; 29 0.5 q in 300 ml distilled H₂O) for 10 min. The 30 coverslips were then dried by placing in a 70°C oven 31 for 45 min. 32

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Colloidal gold suspension was prepared by adding 11 ml 34 35 distilled H₂O and 6 ml Na₂CO₃ (36.5 mM) to 1.8 ml AuHCl₄

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WO 99/54356 PCT/GB99/01211

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1 (14.5 mM). The mixture was heated to 95°C at which

- 2 point 1.8 ml of freshly prepared 0.1% formaldehyde
- 3 solution was added; the temperature was maintained at
- 4 95°C. A suspension of colloidal gold was formed which
- 5 was brown to absorbed light and blue to transmitted
- 6 light.

7

- 8 The gold suspension, was then added to petri dishes
- 9 containing individual coverslips and the plates
- incubated at 37°C for 45 min. After washing with CFS (3
- 11 x 4 ml) to remove unattached gold particles, the
- coverslips were transferred to 6-well cluster dishes
- 13 and UV sterilised.

14

- 15 Endothelial cells (SK HEP-1 and BRCE) in culture media
- 16 (0.3 ml) were seeded onto the coverslips at an
- approximate density of 5 x 10³ cells per well. The cells
- were allowed to plate down for 2 h at 37°C after which
- 19 the treatments were added. Assay systems were
- 20 maintained for a further 18 h after which the cells
- were fixed using 3% gluteraldehyde in cacodylate buffer
- 22 (pH 7.2).

23

- The assays were examined using a Leica DM1RB phase
- 25 contrast microscope and Q500MC image analysis system
- 26 incorporating a JVC TK-1280E colour camera (Leica,
- 27 Milton Keynes, UK). The track images of at least 30
- 28 cells were video-captured and the area (representing
- 29 migration response) determined for each. Statistical
- 30 analysis of these areas was then carried out using
- 31 Macintosh Instat software to perform both Kruskal-
- 32 Wallis analysis of variance and Mann-Whitney U-tests in
- order to compare the treatment groups with controls.

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35 RESULTS

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1 Proliferative response

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- 3 All peptides were examined for their ability to
- 4 influence the growth of T47-D and SK-Hep 1 cell lines.
- 5 At concentrations of peptide up to $100\mu M$, no
- 6 significant effects were observed in either cell line.

7 8

Mechanism of action

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10 It had shown previously that mEGF₍₃₃₋₄₂₎ could inhibit the

11 EGF-stimulated angiogenic response in the early chick

as well as blocking the basal and EGF-stimulated

motility of primary and established endothelial cells.

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During the present study it is shown that mEGF (33-42) also

inhibits the angiogenic effects of laminin (Nelson et

al 1995). Furthermore, it is demonstrated that the

anti-angiogenic effects of mEGF(33-42) are mediated solely

19 through the high affinity 67 kDa laminin receptor (67-

20 LR) and not through the EGF receptor.

21

The study also confirms that $mEGF_{(33-42)}$, Lam. β -1₍₉₂₅₋₉₃₃₎ and

23 laminin are equipotent in 125I-laminin displacement

24 receptor assays, and that both of the small peptidal

25 ligands have similar potencies in specific laminin cell

26 attachment assays.

27

In addition, it is shown that the commonly used chick

29 angiogenesis models are not appropriate to the study of

laminin mediated human angiogenesis: although it is

31 confirmed that Lam. β -1₍₉₂₅₋₉₃₃₎ acts as a partial laminin

32 antagonist in chick, it was found to be a pure agonist

in mammalian cell lines. This is a highly significant

34 point given that pharmaceutical companies (such as

35 Angiotech, Vancouver, BC) are using the chick CAM assay

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PCT/GB99/01211 WO 99/54356

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as the sole screening method for the discovery of anti-1 angiogenic lead compounds. This may be inappropriate 2

for use in human disease. 3

4

This study is the first to show that the YIGSR-receptor 5 is, in fact, the 67 kda high affinity laminin receptor 6 (67-LR). In collaboration with Professor Archer's team 7. at the Department of Ophthalmology, Royal Victoria 8 Hospital, Belfast, it has been determined that the 67-9 LR is preferentially expressed in new vessels during 10

oxygen-induced retinopathy in neonatal mice. 11

12 13

Peptide antagonist development

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The N-terminus of Lam. β -1₍₉₂₅₋₉₃₃₎ is not necessary for receptor recognition and the agonist activity of YIGSR peptide (Ostheimer et al 1992, Kawasaki et al 1994).

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However, alanine scanning of the starting peptide 19 $(mEGF_{(3)-42})$ indicated that residues at positions 1, 2, 20 3, and 6 (peptidesVI, VII, VIII and X respectively), 21 are essential for receptor mediated activities as 22 determined by 125I-laminin displacement and cell 23 attachment to laminin through the 67-LR. Substitution 24 of these individual residues by alanine leads to a 25 dramatic decrease in receptor affinity observed as an 26 increased IC50 (Table 2) and a parallel decrease in 27 their ability to block adhesion to laminin (increased 28 EC₅₀; Table 2). Characterisation of these analogues with 29 regard to effects on motility, largely confirmed these 30 findings although there was one exception; peptide 31 VIII. Results from the migration assay identified this 32 sequence (alanine for cysteine (P1)) as being a weak 33 laminin agonist despite there being a much reduced 34 response in the other two assays. It is suggested that

24

this peptide may influence laminin receptor mediated

2 migration through an alternative mechanism (Scott

3 1997).

4

5 Substitution at P10 (alanine for cysteine (peptide X)

6 retains both receptor binding and adhesion displacing

7 activities but has the effect of changing the

8 antagonistic parent into an agonist analogue. This

9 reflects the response the agonism of Lam. β -1₍₉₂₅₋₉₃₃₎,

which also lacks the C-terminal cysteine, and suggests

that this cysteine is not essential for receptor

recognition, but is required for antagonism of mEGF(33.

13 42).

14

15 Studies have reported that the positive charge offered

16 by arginine (P9) is essential for the biological

activity of Lam. β -1₍₉₂₅₋₉₃₃₎ (McKelvey et al 1991, Kawasaki

et al 1994). Glutamate substitution for arginine

19 generates a negative charge at this position with

20 corresponding loss of biological activities (Kawasaki

21 et al 1994).

22

23 However, the substitution of arginine (P9) with

24 positively-charged lysine (McKelvey et al 1991) also

25 results in complete loss of ligand binding and

26 biological activities, suggesting that the mere

27 presence of a positive charge at this position is, in

itself, insufficient for receptor recognition. This

29 modelling studies suggest that H-bonding of the

30 quanidino group of the arginyl residue to the aromatic

31 sidechain of the tyrosyl residue (P5) in the consensus

32 sequence GYXGXR presents an acceptable motif for 67-LR

activation by both mEGF₍₃₃₋₄₂₎ and Lam. β -1₍₉₂₅₋₉₃₃₎.

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35 Substitution of tyrosine (P5) with a conformationally

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restricted mimetic (tetrahydroisoquinoline-3-carboxylic 1 acid; Tic-OH) in peptide V converted the antagonist 2 mEGF(1).42) into an agonist. This residue substitution 3 generates a predicted conformation unlikely to be able 4 to form H-bonds. Although both receptor binding and 5 adhesion responses were retained in this peptide the 6 loss of antagonism would suggest that H-bonding between 7 tyrosine (P5) and the arginine (P9) is important for 8 these antagonist activities. 9 10

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Modelling studies suggested that citrulline (an uncharged arginine mimetic) would also be capable of forming this H-bonded motif.

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It was found that replacement of arginine (P9) with citrulline (peptide IV) increased both receptor binding and inhibition of attachment to laminin substrata whilst retaining antagonist migratory response (Table 2), reinforcing the observation that it is not the positive charge that is required rather than an active conformation generated by hydrogen bonding. These findings thus identify H-bonding between P5 and P9 as being more important than the charge at the P9 arginine in determining antagonist activity.

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Subsequent strategies involved the substitution of variant residues in the antagonistic $\text{mEGF}_{(33-42)}$ with those present in the agonistic $\text{Lam.}\beta\text{--}1_{(925-933)}$ sequence (peptides I-III), in an effort to identify key amino acids in the *C*-terminal regions (P5-10) of the two ligands responsible for their contrasting bioactivities.

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34 Substitution of isoleucine (P6) for serine (peptide I) 35 resulted in both reduced receptor affinity and potency

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611

WO 99/54356 PCT/GB99/01211

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in displacement of cell adhesion to laminin. However, this analogue retained weak antagonist activities in 2 the motility assay. It is therefore of interest that 3 studies on the YIGSR sequence indicate that residue 4 substitution, at the position taken by isoleucine in 5 the pentapeptide, are well tolerated and may increase 6 potency (Kawasaki et al 1994). 7 8 Replacement of aspartate (P8) with serine (peptide II) 9 resulted in a complete loss of biological function. as 10 did peptide III encompassing both the former 11 (isoleucine (P6) for serine) and latter (serine (P8) 12 for aspartate) substitutions. Since this mEGF(33-42) 13 analogue sequence (peptide II) encompasses the active 14 YIGSR amino acid sequence agonist, it is suggested that 15 this loss of activity may be attributed to the valine 16 (P2) and isoleucine (P3) residues in the N-terminal 17 half of $mEGF_{(33-42)}$. Alternatively, addition of a C-18 terminal cysteine to the YIGSR sequence is known to 19 reduce potency (Kawasaki et al 1994). Additional 20 peptides incorporating the valine (P2) and isoleucine 21 (P3) substitutions are currently under investigation. 22 23 The determination of the minimum core peptide structure 24 is ongoing and involves similar characterisation 25 studies on a number of sequences truncated at the C-26 27 terminal. 28 These studies have thus identified an important 29 antagonist of 67-LR mediated activities in peptide IV. 30 The sequence, (AcC(Acm)-VIGYSGD-[Cit]-C-(Acm)-NH2.), may 31 provide an important template for anti-angiogenic drugs 32 in that it is resistant to cleavage by trypsin-like 33

proteases and has been identified as being more potent

than mEGF(33-42) in screening procedures.



PCT/GB99/01211

WO 99/54356

1	<u>Advantages</u>
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The advantages of the invention, and the ways in which disadvantages of previously known arrangements are overcome include:

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1. Unlike the native 67LR ligand (laminin β -l_{925.933}), which is angiogenic in human models, the mEGF₃₃₋₄₂-derived agents are anti-angiogenic in human models.

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12 2. mEGF₃₃₋₄₂ has the advantage of inhibiting both 13 laminin- and EGF-stimulated angiogenesis.

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3. mEGF₃₃₋₄₂ prevents tumour cell attachment to
 basement membranes.

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